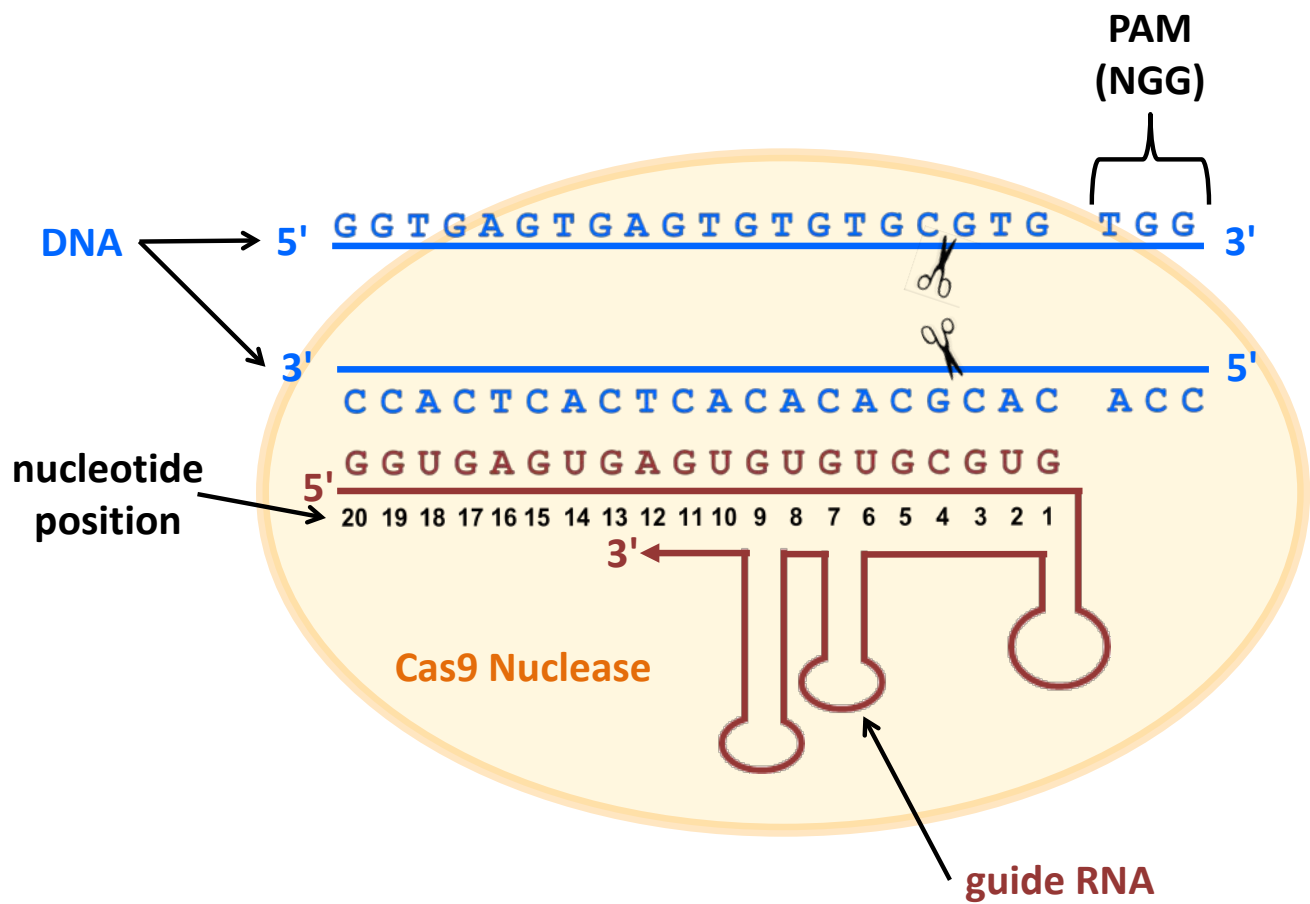


Supplementary Results

Titration of sgRNA- and Cas9-expressing plasmid amounts used for the EGFP disruption assay

Single guide RNAs (sgRNAs) were generated for three different sequences (**Supplementary Fig. 2a**) located upstream of *EGFP* nucleotide 502, a position at which the introduction of frameshift mutations via non-homologous end-joining can robustly disrupt expression of EGFP^{1, 2}. For each of the three target sites, we initially transfected a range of sgRNA-expressing plasmid amounts (12.5 to 250 ng) together with 750 ng of a plasmid expressing a codon-optimized version of the Cas9 nuclease into our U2OS.EGFP reporter cells bearing a single copy, constitutively expressed *EGFP-PEST* reporter gene. All three RGENs efficiently disrupted EGFP expression at the highest concentration of sgRNA plasmid (250 ng) (**Supplementary Fig. 2b**). However, RGENs for target sites #1 and #3 exhibited equivalent levels of disruption when lower amounts of sgRNA-expressing plasmid were transfected whereas RGEN activity at target site #2 dropped immediately when the amount of sgRNA-expressing plasmid transfected was decreased (**Supplementary Fig. 2b**). We next titrated the amount of Cas9-encoding plasmid (range from 50 ng to 750 ng) transfected into our U2OS.EGFP reporter cells and assayed for EGFP disruption. As shown in **Supplementary Fig. 2c**, target site #1 tolerated a three-fold decrease in the amount of Cas9-encoding plasmid transfected without substantial loss of EGFP disruption activity. However, the activities of RGENs targeting target sites #2 and #3 decreased immediately with a three-fold reduction in the amount of Cas9 plasmid transfected (**Supplementary Fig. 2c**). Based on these results, we used 25ng/250ng, 250ng/750ng, and 200ng/750ng of sgRNA-/Cas9-expressing plasmids for *EGFP* target sites #1, #2, and #3, respectively, for the experiments described in the main text.

We do not understand the reasons why some sgRNA/Cas9 combinations work better than others in disrupting EGFP expression nor do we know why some of these combinations are more or less sensitive to the amount of plasmids used for transfection. Although it is possible that the range of off-target sites present in the genome for these three sgRNAs might influence each of their activities, we did not observe any differences in the numbers of genomic sites that differ by one to six bps for each of these particular target sites (**Supplementary Table 3**) that would account for the differential behavior of the three sgRNAs.

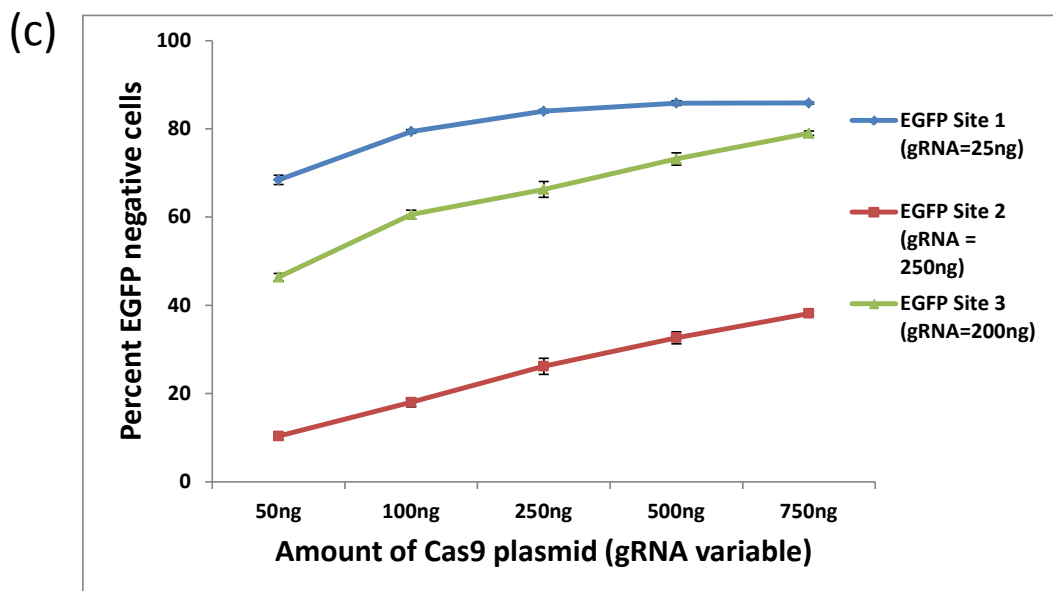
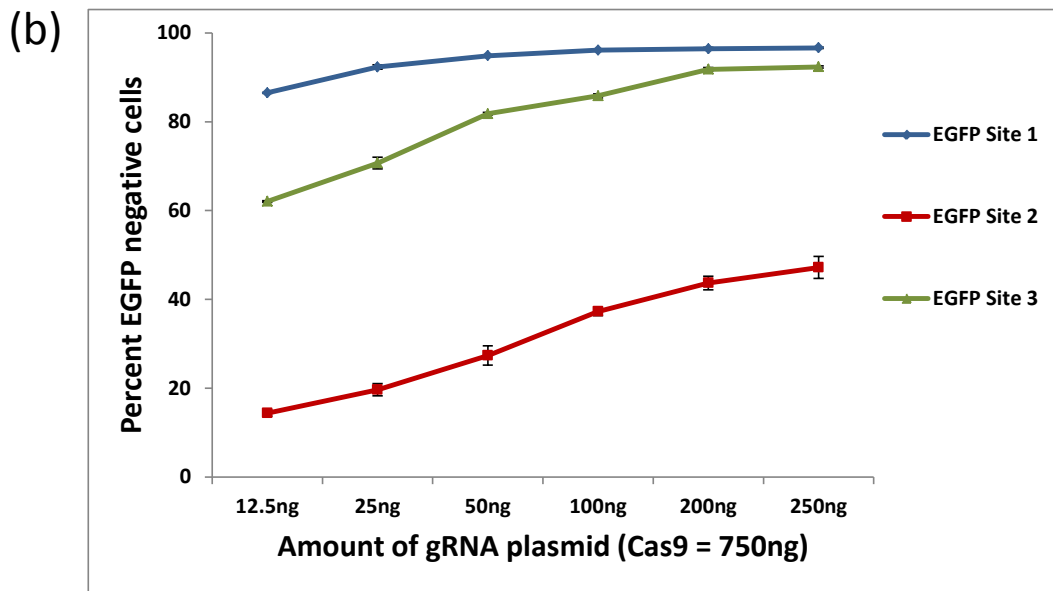


Supplementary Fig. 1 Schematic illustrating a sgRNA/Cas9 nuclease complex bound to its target DNA site

Scissors indicate approximate cleavage points of the Cas9 nuclease on the genomic DNA target site.

Note the numbering of nucleotides on the guide RNA proceeds in an inverse fashion from 3' to 5'.

(a) EGFP Site 1 GGGCACGGGCAGCTTGCCGGTGG
 EGFP Site 2 GATGCCGTTCTTCTGCTTGTTCGG
 EGFP Site 3 GGTGGTGCAGATGAACATTCAAGGG



Supplementary Fig. 2 Titration of sgRNA- and Cas9-expressing plasmid amounts used for the human cell-based EGFP disruption assay

(a) Sequences for three RGEN target sites in *EGFP* used in this study. (b) Activities of RGENs in the EGFP disruption assay performed using varying amounts of sgRNA-expressing plasmids. (c) Activities of RGENs in the EGFP disruption assay performed using varying amounts of Cas9-expression plasmid.

Target 1 (VEGFA Site 1):

OT1-3

AGACAGGACATTCTGACACCCCAGGAGCAAACTCCCTCCATCCCACAAATCCGTCCTAGATGTGCA Wild-type x18
AGACAGGACATTCTGACACCCCCATCCCACAAATCCGTCCTAGATGTGCA Δ53
AGACAGGACATTCTGACACCCCACAAAATCCGTCCTAGATGTGCA Δ17
AGACAGGACATTCTGACACCCCACAAAATCCGTCCTAGATGTGCA Δ15
AGACAGGACATTCTGACACCCCACAAAATCCGTCCTAGATGTGCA Δ2

AGACAGGACATTCTGACACCCCAGGAGGCAAACTCCCTCCATCCCACAAATCCGTCCTAGATGTGCA +2

OT1-6

GAGAGAGGCTCCCCATCACGGGGGAGGGAGTTTGCTCCTGGGAACCTGTGATCCCCACAGGGAAACA Wild-type x87
GAGAGAGGCTCCCCATCACGGGGGGAGGGAAACA Δ35 x3
GAGAGAGGCTCCCCATCACGGGGGGAGGGAAACA Δ14 x1

OT1-11

TGGACTCTACCCACTGAATGCAGGAGCAAACTCCCTCCCCGAGTTGTGACAGCAAAATGTCTC Wild-type x27
TGGACTCTACCCACTGGCAGGAGCAAACTCCCTCCCCGAGTTGTGACAGCAAAATGTCTC Δ43
TGGACTCTAGCAGGAGCAAACTCCCTCCCCGAGTTGTGACAGCAAAATGTCTC Δ25
TGGACTCTACGCAGGAGCAAACTCCCTCCCCGAGTTGTGACAGCAAAATGTCTC Δ24
TGGACTCTACCCACTGAATGCAGGAGCAAACTCCCTCCCCGAGTTGTGACAGCAAAATGTCTC Δ15
TGGACTCTACCCACTGAATGCAGGAGCAAACTCCCTCCCCGAGTTGTGACAGCAAAATGTCTC Δ15
TGGACTCTACCCACTGAATGCAGGAGCAAACTCCCTCCCCGAGTTGTGACAGCAAAATGTCTC Δ7
TGGACTCTACCCACTGAATGCCAGGATTCCCTCCCCGAGTTGTGACAGCAAAATGTCTC Δ6
TGGACTCTACCCACTGAATGCCCTGGAGTTGTGACAGCAAAATGTCTC Δ6
TGGACTCTACCCACTGAATGCCAGΔ3
TGGACTCTACCCACTGAATGCCAGΔ3

TGGACTCTACCCACTGAATGCCAGGATGGAAGATAATTTTTCCATAGACCAGGGTGGGGGAATGGTTGGGATGATTCAAGCACATCAC
ATTTATTGTGCACTTTATTCTATTACTATTATATTGTAATGTACTAAAAAAATTATACAACTACCATAATGTAGAACCAGTGGGAG
CGCAAACTCCCTCCCCGAGTTGTGACAGCAAAATGTCTCA +158

TGGACTCTACCCACTGAATGCCAGGATGGAAGATAATTTTTCCATAGACCAGGGTGGGGGAATGGTTGGGATGATTCAAGCACATCAC
AGACATTGCCAAATGTCCCT +23 (Δ2 +25)

Target 2 (VEGFA Site 2):

OT2-2

ACCCACCTCCCTATCCTCAAAACTTGGCAGAGGCGGGTGGAGGGGCCCTAGGAGCGCCTTGGT Wild-type x30
ACCCACGCAGAGGCGGGTGGAGGGGCCCTAGGAGCGCCTTGGT Δ51
ACCCACCTCCGCAGAGGCGGGTGGAGGGGCCCTAGGAGCGCCTTGGT Δ36
ACCCACCTCCCTATCCTCAAAACTTGCCAGAGGGGTGGAGGGGCCCTAGGAGCGCCTTGGT Δ33
ACCCACCTCCCTATCCTCAAAACTTGCCAGAGGGGTGGAGGGGCCCTAGGAGCGCCTTGGT Δ18
ACCCACCTCCCTATCCTCAAAACTTGCCAGAGGGGTGGAGGGGCCCTAGGAGCGCCTTGGT Δ10
ACCCACCTCCCTATCCTCAAAACTTGCCAGAGGGGTGGAGGGGCCCTAGGAGCGCCTTGGT Δ7
ACCCACCTCCCTATCCTCAAAACTTGCCAGAGGGGTGGAGGGGCCCTAGGAGCGCCTTGGT Δ6

ACCCACCTCCCTATCCTCAAAACTGCCAGAGGGGTGGAGGGGCCCTAGGAGCGCCTTGGT
CTGGCAGTCAGTGCGTTATCTGTCACACTTCTACAAGGGGCTCCCTGCATTCTGA +21 (Δ40, +61)

ACCCACCTCCCTATCCTCAAAACTTGCCAGAGGGGTGGAGGGGCCCTAGGAGCGCCTTGGT
CTGGGACCATAGCCGGGGACCCAGACTCTGGCCACGCTATTCCACCGCGGGGTGGAGGGGCCCTAGGAGCGCCTTGGT +108

OT2-15

TGACTGTCGGTGCCCCACATGTGGCAGATGCCCAGAGGCGGGTGTGGGGGTACTTGTGGGCGT Wild-type x71
TGACTGTCGCGGTGCCCCACATGTGGCAGATGCGGGTGGGGGTACTTGTGGGCGT Δ33
TGACTGTCGGTGCCCCACATGTGGCAGATGCGGGTGGGGGTACTTGTGGGCGT Δ23
TGACTGTCGGTGCCCCACATGTGGCAGAGTGGGTGGGGGTACTTGTGGGCGT Δ15
TGACTGTCGGTGCCCCACATGTGGCAGAGTGGGTGGGGGTACTTGTGGGCGT Δ14
TGACTGTCGGTGCCCCACATGTGGCAGATGCGGGTGGGGGTACTTGTGGGCGT Δ4
TGACTGTCGGTGCCCCACATGTGGCAGATGCGGGTGGGGGTACTTGTGGGCGT Δ4
TGACTGTCGGTGCCCCACATGTGGCAGATGCGGGTGGGGGTACTTGTGGGCGT Δ2

TGACTGTGGTCCCCACATGTGGCAGATGCCAGAG_{TT}GCAGGTGTGGGGGTACTTGTCGGCGTT +2
TGACTGTGGTCCCCACATGTGGCAGATGCCAGAG_{CT}GCGGGGTGTGGGGGTACTTGTCGGCGTT +2

OT2-24

ACAAGATGACTATGTCCCTCTGGGCCCATCCTCCC_{CTCCCCACCCACCCGCCCTCAGG}CTTGAAGA Wild-type x8
ACAAGA₋₋₋_> Δ121
<₋₋₋ CTCAGGCTTGAAGA Δ82
ACAAGATG₋₋₋_> Δ80
<₋₋₋_> Δ79
ACAAGATGACTATGTCCCTCTGGGCCCATCCTCC₋₋₋_> Δ42
ACAAGATGACTATGTCCCTCTGGGCC₋₋₋_> CTTGAAGA Δ34
ACAAGATGACTATGTCCCTCTGGGCCCATCCTCC₋₋₋_> CAGGCTTGAAGA Δ14
ACAAGATGACTATGTCCCTCTGGGCCCATCCTCC₋₋₋_> CCTCAGGCTTGAAGA Δ12
ACAAGATGACTATGTCCCTCTGGGCCCATCCTCC₋₋₋_> GCAGGCTTGAAGA Δ11
ACAAGATGACTATGTCCCTCTGGGCCCATCCTCC₋₋₋_> CTTGAAGA Δ11
ACAAGATGACTATGTCCCTCTGGGCCCATCCTCC₋₋₋_> GCTCAGGCTTGAAGA Δ11

ACAAGATGACTATGTCCCTCTGGGCC_{GCCTCAAGTGATCCAGCTGCC}CTGGCC_{CAA}AAGTGCTAGCAGTACAGATGTGAGC_{CATG}CCTGGC_{CATT}
{CCTGGC}TATTGCAACATCCC{AT}CTGTGAAGCAGGTTCTGCAAGTGACAGCAAGAAGAGCACAGGGCCAAAAAAACTTGTCC_{TTA}
GAAAGGATCTACCTTTAGGCTGAGAATGGCA +76 (Δ43 +119)

Target 3 (VEGFA Site 3):

OT3-2

GAGTGAGAGAGCGAGTGAGTGAGTGAGTGAGTGAGTGA₋₋₋₋GTGTGAGTGAGTGAGTGAGTGA Δ4
GAGTGAGAGAGCGAGTGAGTGAGTGAGTGAGTGA₋₋₋₋GTGTGAGTGAGTGAGTGAGTGA Δ6 x2

OT3-9

GTGTTGGATGCGGGAGTGAGTGAGTGAGTGCGTGC_{GGGTGG}CGATGCAAGCGTGTGCGAATGCGTG x173
GTGTTGGATGCGGG₋₋₋_> Δ80
GTGTTGGATGC₋₋₋_> GCGTG Δ50
GTGTTGGATGCGGGAGTGGTGAGTGAGTGA₋₋₋₋GTGGCGATGCAAGCGTGTGCGAATGCGTG Δ10

GTGTTGGATGCGGGAGTGGTGAGTGAGTG_{AA}GTGCGGGTGCGATGCAAGCGTGTGCGAATGCGTG +2

OT3-18

TTTCAAAGACAGTAGATCTAAATGT_{CCTCACGCACACACTCACCCACACATAAAAGGTGGTA}CTG Wild-type x27
TTTCAAAGACAGTAGATCT_{AAATGT}₋₋₋₋TAAAGGTGGTAACTG Δ32
TTTCAAAGACAGTAGATCTAAATGT₋₋₋CATAAAAGGTGGTAACTG Δ23
TTTCAAAGACAGTAGATCTAAATGT_{CCT}₋₋₋₋CACATAAAAGGTGGTAACTG Δ18 x4
TTTCAAAGACAGTAGATCTAAATGT_{CCT}₋₋₋₋CCACACATAAAAGGTGGTAACTG Δ15
TTTCAAAGACAGTAGATCTAAATGT_{CCT}₋₋₋₋CTCACCCACACATAAAAGGTGGTAACTG Δ12
TTTCAAAGACAGTAGATCTAAATGT_{CCTCA}₋₋₋₋CACACACTCACCCACACATAAAAGGTGGTAACTG Δ2

TTTCAAAGACAGTAGATCTAAATGT_{CACAGGCTGGAGTACAGTGGCATGATATCAGCTCA}TGCAATCTGGGCTCCGGTTCAAG
CCATGACACACTCACCCACACATAAAAGGTGGTAAC +63

Target 4 (EMX1):

OT4-1

ACCTGTACATCTGCACAAGATTGCC_{TTTCTTCTTCTGCTCTAAC}CTGACAATC Wild-type x20
ACCTGTACATCTGCACAAGATTGCC_{TTTACTCC}₋₋₋₋₋ATC Δ64
ACCTGTACATCTGCACAAGATTGCC_{TTTACTCC}₋₋₋₋₋ACAATC Δ28
ACCTGTACATCTGCACAAGATTGCC_{TTTACTCC}₋₋₋₋₋ACTCTGACAATC Δ20
ACCTGTACATCTGCACAAGATTGCC_{TTTACTCC}₋₋₋₋₋CTTCTGCTCTAACCTGACAATC Δ20
ACCTGTACATCTGCACAAGATTGCC_{TTTACTCC}₋₋₋₋₋CAATC Δ19
ACCTGTACATCTGCACAAGATTGCC_{TTTACTCC}₋₋₋₋₋TGCTCTAACCTGACAATC Δ14
TCCTGTACATCTGCACAAGATTGCC_{TTTACTCC}₋₋₋₋₋CTTCTTCTGCTCTAACCTGACAATC Δ8

Supplementary Fig. 3 Sequences of off-target indel mutations induced by RGENs in human U2OS.EGFP cells

Wild-type genomic off-target sites recognized by RGENs (including the PAM sequence) are highlighted in yellow and numbered as in **Table 1** and **Supplementary Table 2**. Note that the complementary strand is shown for some sites. Deleted bases are shown as red dashes on a grey background. Inserted bases are italicized and highlighted in blue.

Target 1 (VEGFA Site 1):

OT1-3

TCAGACAGGACATTCTGACACC **CCAGGGAGCAA**ACTCCTCCATCCCACAAATCCGTCTTAGATGTG Wild-type x41

TCAGACAGGACATT----- **CAA**ACTCCTCCATCCCACAAATCCGTCTTAGATGTG Δ15

TCAGACAGGACATTCT----- **GAG**CAA^{ACTCCTCCATCCCACAAATCCGTCTTAGATGTG} Δ10

TCAGACAGGACATTCTGAC----- **GCA**AA^{ACTCCTCCATCCCACAAATCCGTCTTAGATGTG} Δ9

TCAGACAGGACATTCTGACAC----- **GCA**AA^{ACTCCTCCATCCCACAAATCCGTCTTAGATGTG} Δ7

TCAGACAGGACATTCTGACACCCCCAG----- **GCA**AA^{ACTCCTCCATCCCACAAATCCGTCTTAGATGTG} Δ2 x6

TCAGACAGGACATTCTGACACCCCCAGGA **TG**TCCCTCCCTCCATCCCACAAATCCGTCTTAGATGTG +2

TCAGNCAGGACATTNNNGNCACCCCAGGA **AAACNN**GAGTTTCGNTNCNNNGANNGTCAGACCCAGNAGCAA^{ACTCCTCCATCCCACAAATC}CGTCTTAGATGTG +38

TCAGACAGGACATTNTGACACCCCCAGGA **GTNTGCACNT**CAGTTTCTTTANTATGTNGNNNNGGGCANGNACAAANNTTN **GCAA**ACTCC **CTCC**ATCCCACAAATCCGTCTTAGATGTG +54

TCAGACAGGACATTCTGACACCCCCAGGA **ATGTTTGT**TTGAGTCAGAGTCTCTCTTTGTCA^{CC}CAGGCTGGAGTGCA^{GT}GGAA^{AC}CTGTGCC **TTT**GTATATCCTCTTGAAGGTTAAAGAGTCATCATGGATCANCNNATAAAGCAA^{ACTCC}NTCCATCC +116

TCAGACAGGACATTCTGACACCCCCAGGA **TAAGCGGCAGGGTCGGA**ACAGGAGAGCGCACGAGGGAGCTCAGGGGGAA^{AC}GCCTGGTATC **TTT**ATAGCCTGTCGGGTTTCGCCACCTCTAGGGAGTTCTGAGCTCTGACGCCGGCGTGTAGGAGATAGCAGGCCGTTAATGACCATC **CCAGCCGAA**TTCTCACTGTGCA^{GT}AGT^{AG}GACTCAGGGAGGCTGAGTGTCCCAGGC^{CT}GTG^{CC}CAGATGAGGCCACGCTGAGACT **GTG**CAA^{ACT}CCCTCATCCCACAAATCCGTCTTAGATGTG +247

OT1-6

TGGAGAGAGGGCTCCATC **ACGGGGGAGGGAGTTG**CTCCTGGGAACCTGTGATCCCCACAGGGAAC Wild-type x88

TGGAGAGAGGGCTCCATCAGGGGGAGGGAGTTG----- **CCT**GTGATCCCCACAGGGAAC Δ11

TGGAGAGAGGGCTCCATCAGGGGGAGGGAGTTG----- **GGG**AACCTGTGATCCCCACAGGGAAC Δ7

OT1-11

AGCATCGCTGGACTCTACCCACTGAATGCCAGG **AGTC**CAGACGATTGAATGTATCAACTTGGCACATTGCCATCAACTGGTAGTGCTCA

AAAATATCCATTGCTGTGATCAGTAATGCCACAGGGTGACCATTAAAGGACAGAGTCCATTTTATCCATCCTT **AGCAA**ACTCCCCTCC

CCGAGTTGTGACAGCAA +133

AGCATCGCTGGACTCTACCCACTGAATGCCAGG **AGCC**CTTCCCTCCTCCCTCTCTCCAGAGGTCTGCCAGATCAGGTTGGAGGTCCCT **CTTT**GTCTTATGCCATTCTCCCCAGGC^{ACT}TGGAGGAGGGACTGTTAGTGTCAAGTCTTCTGT^{TT}ACTGTTGGCAAC **TT**CCCTCTCCCGAGTTGTGACAGCAA +142

AGCATCGCTGGACTCTACCCACTGAATGCCAGG **TGTTTGT**TTGCGACGGANTCTCACTCTGTCGNCCGGCTGGAGTGCA^{AN}GGCACAN **TT**CCCTC **AN**CTGACTGCNATGTCCGCCTCCGGATTCAAGT^GATTCTCTGCCAGCCTCCGAGTAGCTGGATTATAGGTGCC^{CT}GCCAC **CAT**GCC^{CT}GGCTAATT^{TT}TTTTTTTAAATGGAGTCTCACTCTGTTGCCCGAGTTGTGACAGCAA +186

Target 2 (VEGFA Site 2):

OT2-2

CACCTCCCTATCCTCAAAACTTGG **CCAGAGGCGGGGTGGAGGGGGCC**CTAGGAGCGCCTTGGTGGGA

CACCTCCCTATCCTCAAAACTTGGCCAGAGGCGGGGT----- **GGAGCGCCTTGGTGGGA** Δ13

CACCTCCCTATCCTCAAAACTT----- **GGCGGGGTGGAGGGGGCC**CTAGGAGCGCCTTGGTGGGA Δ7

CACCTCCCTATCCTCAAAACTT----- **GACGGGGTGGAGGGGGCC**CTAGGAGCGCCTTGGTGGGA Δ6

CACCTCCCTATCCTCAAAACTTGGC----- **AGGCGGGGTGGAGGGGGCC**CTAGGAGCGCCTTGGTGGGA Δ2

CACCTCCCTATCCTCAAAACTTGGCCAGAGACTTANACCTAANACCTCAA^{ACT}ATGAGACTGCTACNAGAGAACATCANAAA^{ACT}TTCCA **GGAC**ATTCTCTGNGGGTGGAGGGCCCTANGAGCGCCTTGNNGGA +74

CACCTCCCTATCCTCAAAACTTGGCCAGAG**TCTACAG**ATT^TAAAATATTACCA^GT^TAATCATGACACATATTGTTATT^TCAA^AATT^TTTCTAGTTAAACCCACCATT^TATAACCA^{TT}ATT^TGATATT^TAAAATT^TGTATTAAACACCCACCA^AATT^TTACAGC

GGGGTGGAGGGGCCCTAGGAGCGCCTGGTGGGA +150

OT2-15

GTCGGTCCCCACATGTGGCAGATGC **CCAGAGGGGGGTGTGGGGGTACTTTGTGGCGTTTGGG** Wild-type x79
GTCGGTCCCCAC-----GCGGGGTGTGGGGGGTACTTTGTGGCGTTTGGG Δ19
GTCGGTCCCCACATGTGGCAGATGC **CCAGAGGGGGGTACTTTGTGGCGTTTGGG** Δ12
GTCGGTCCCCACATGTGGCAGATGCCAGA-----GGTGTGGGGGTACTTTGTGGCGTTTGGG Δ5
GTCGGTCCCCACATGTGGCAGATGCCCA-----GGGTGTGGGGGTACTTTGTGGCGTTTGGG Δ5
GTCGGTCCCCACATGTGGCAGATGCCAGA-----GGGTGTGGGGGTACTTTGTGGCGTTTGGG Δ4
GTCGGTCCCCACATGTGGCAGATGCCCA-----GGGGGTGTGGGGGTACTTTGTGGCGTTTGGG Δ4
GTCGGTCCCCACATGTGGCAGATGCCCA-----AGGCGGGGTGTGGGGGTACTTTGTGGCGTTTGGG Δ2 x2

OT2-24

GTCCCTCTGGGCCCCATCCTCCC **CTCCCCACCCACCCGCCCTCAGGCTTGAAAGAGGAAAGAAAGAGCA**
GTCCCTCTGGGCCCC-----ANAGNANNNNNNNN Δ36
GTCCCTCTGGGCCCCATCCTCCCCTCCCC-----TGAAGAGGAAAGAAAGAGCG Δ19
GTCCCTCTNNNCNNCNT-----CCNCNCCTCAGGCTTGAAAGAGGAAAGAAAGAGCG Δ17
GTCCCTCTGGGCCCCATCCTCCCCTCCTC-----CCTCAGGCTTGAAAGAGGAAAGAAAGAGCG Δ13
GTCCCTCTGGGCCCCATCCTCCCCTCCCC-----CTCAGGCTTGAAAGAGGAAAGAAAGAGCG Δ10
GTCCCTCNNNNNNNCCTCCTCCC-----NNCCNCNCCTCAGGCTTGAAAGAGGAAAGAAAGAGCG Δ8
GTCCCTCTGGGCCCCATCCTCCCCTCCCCACCCA-----CTCAGGCTTGAAAGAGGAAAGAAAGAGCG Δ6
GTCCCTCTGGGCCCCATCCTCCCCTCCCCACCCA-----CCTCAGGCTTGAAAGAGGAAAGAAAGAGCG Δ5

GTCCCTCTGGGCCCCATCCTCCCCTCCCCACCCACCCCG **CATCGTACGTGTCCTTACTAAGCTGCAATTGGCATTTCAAGCTAACGTGAA**
GTTCGA CCTCAGGCTTGAAAGAGGAAAGAAAGAGCG +58

Target 3 (VEGFA Site 3):

OT3-2*

GAGAGCGAGTGAGTGAGTGAGTGAGTGAGTGAGTGAGTGAGTGAGTNNTNNNNATTTACACACATACGAGCCGAAGCATAAAGTGTAAAGCCTAGGGTGCCTA

ATGAGTGAGCTAACTCACATTAAATTGCGTTGCGCTCACTGCCGCTTCCAGTCGGAAACCTGTCGTGCCAGCTGCATTAATGAATCGGC
CAANNNNNGNNNAGAGGCGGTTGCGTATTGGCGCTCTCCGCTTCCTCGCTCACTGACTCGCTGCCCTCGGTGTTGGCTGCGCG
CGGTATCAGCTCACTAAAGGCGGTAATACGGTTATCCACANAANGGACTCGGCTTGTGTCGGTGAATT +280

OT3-9

TGGAGGTGTTGGGATGCGGGAGTG **GGTGAGTGAGTGCGTGCGGGTGGCGATGCAAGCGTGTGCAAT** Wild-type x101

TGGAGGTGTTGGGATGCGGGAGTG -----GTGCGTGCGGGTGGCGATGCAAGCGTGTGCAAT Δ8

TGGAGGTGTTGGGATGCGGGAGTG -----GTGCGGGTGGCGATGCAAGCGTGTGCAAT Δ8

OT3-18

CAAAGACAGTAGATCTTAAATGT **CCTCACCGCACACACTCACCCACACATAAAAGGTGGTAACTGTGT** Wild-type x64

CAAAGACAGTAGATCTTAA-----GCACACACTCACCCACACATAAAAGGTGGTAACTGTGT Δ10

CAAAGACAGTAGATCTTAAATGTCCTCACG -----TCACCCACACATAAAAGGTGGTAACTGTGT Δ7

CAAAGACAGTAGATCTTAAATGTC -----GCACACACTCACCCACACATAAAAGGTGGTAACTGTGT Δ5

CAAAGACAGTAGATCTTAAATGTCCTCACG **CCGACNATTGCATGAAGAATCTGTTAGGCTTTGCGCTGCTCGCGATGTACG**
GGCCAGATAGC ACACACTCACCCACACATAAAAGGTGGTAACTGTGT +71

CAAAGACAGTAGATCTTAAATGTCCTCACG **CAAATTTTATTTGGTTCATGATATGGCTTGGCGTATGCTTTCATTTGAAAATTGCT**
GTTCTTTGACAATTAAAGTGA CTGTTCATGACTACAAGTTGAAAATAAAATTAAATTAAAGAAAAAAATTCCAATGACTGTGCTGTGG
TTGGGCACACACTCACCCACACATAAAAGGTGGTAACTGTGT +157

CAAAGACAGTAGATCTTAAATGTCCTCACG **GTGGAAACATAGTAGATGAGGTGGCATATCATGAAAAGTACCCACGATTATCACCTCANAA**
AAAAGCTAGTTGACTCAACTGATAAAGCGGACCTGAGGTTAATCTACTTGGCTTGGCCATATGATAAAGTTCCGTGGGCACTTCTCA
TTGAGGGTGATCTAAATCCGGACAACTCGGATGTCGAGCACACACTCACCCACACATAAAAGGTGGTAACTGTGT +190

CAAAGACAGTAGATCTTAAATGTCCTCACG -----
TTATTTAGAGACAGAGTCTCACCTGTTGCCAGGCTGGGGTGCAGTGGTACGAACCTGGCTCACTGCAACCTCCGTCTCCTGGGCTCAAG
TGATTATCCTGCCCTAGCCTCCCGAGTAGCTGGGATTACAGGTGCCACCACCCGGCTAATTGGTATTTCACTGAGAGCTGGG
TTTCACCATGTTGCCAGCCTGTTCTCGCACACACTCACCCACACATAAAAGGTGGTAACTGTGT +211 (Δ16 +227)

Target 4 (*EMX1*):

OT4-1

GATTGCCTTTACTCCATG**C**CTTCTTCTTGCTCTA**A**CTCTGACAATCTGTCTTGC**C**CATGCCATAA Wild-type x74
GATTGCCTTTACTC**T**CTTCTGCTCTA**A**CTCTGACAATCTGTCTTGC**C**CATGCCATAA Δ9
GATTGCCTTTACTCCATGC**T**CTTCTGCTCTA**A**CTCTGACAATCTGTCTTGC**C**CATGCCATAA Δ6 x2
GATTGCCTTTACTCCATGC**T**CTTCTGCTCTA**A**CTCTGACAATCTGTCTTGC**C**CATGCCATAA Δ6
GATTGCCTTTACTCCATGC**T**CTTCTGCTCTA**A**CTCTGACAATCTGTCTTGC**C**CATGCCATAA Δ3 x3

Supplementary Fig. 4 Sequences of off-target indel mutations induced by RGENs in human HEK293 cells

Wild-type genomic off-target sites recognized by RGENs (including the PAM sequence) are highlighted in yellow and numbered as in **Table 1** and **Supplementary Table 2**. Note that the complementary strand is shown for some sites. Deleted bases are shown as red dashes on a grey background. Inserted bases are italicized and highlighted in blue. *Yielded a large number of single bp indels

Supplementary Table 1 Sequences of oligonucleotides used to generate expression plasmids encoding sgRNAs/variant sgRNAs targeted to sites in the *EGFP* reporter gene and sgRNAs targeted to six endogenous human gene targets

See accompanying file

Supplementary Table 2 Sequences and characteristics of genomic on- and off-target sites for six RGENs targeted to endogenous human genes and primers and PCR conditions used to amplify these sites

See accompanying file

Supplementary Table 3 Numbers of off-target sites in the human genome for six RGENs targeted to endogenous human genes and three RGENs targeted to the *EGFP* reporter gene

Target Site	Number of mismatches to on-target site						
	0	1	2	3	4	5	6
Target 1 (<i>VEGFA</i> Site 1)	1	1	4	32	280	2175	13873
Target 2 (<i>VEGFA</i> Site 2)	1	0	2	35	443	3889	17398
Target 3 (<i>VEGFA</i> Site 3)	1	1	17	377	6028	13398	35517
Target 4 (<i>EMX</i>)	1	0	1	18	276	2309	15731
Target 5 (<i>RNF2</i>)	1	0	0	6	116	976	7443
Target 6 (<i>FANCF</i>)	1	0	1	18	271	1467	9551
EGFP Target Site #1	0	0	3	10	156	1365	9755
EGFP Target Site #2	0	0	0	11	96	974	7353
EGFP Target Site #3	0	0	1	14	165	1439	10361

Off-target sites for each of the six RGENs targeted to the *VEGFA*, *RNF2*, *FANCF*, and *EMX1* genes and the three RGENs targeted to EGFP Target Sites #1, #2 and #3 were identified in human genome sequence build GRCh37. Mismatches were only allowed for the 20 nt region to which the sgRNA anneals and not to the PAM sequence.

Supplementary Table 4 Indel mutation frequencies at on- and off-target genomic sites induced by different amounts of Cas9- and sgRNA-expressing plasmids for the RGEN targeted to *VEGFA* Target Site 3

Site	Sequence	250ng sgRNA/750 ng Cas9 Mean indel frequency (%) ± SEM	12.5ng sgRNA/250 ng Cas9 Mean indel frequency (%) ± SEM
T3 (On-target)	GGTGAGTGAGTGTGTGCGGTG	49.4 ± 3.8	33.0 ± 3.7
OT3-1	GGTGAGTGAGTGTGT <u>T</u> GTGAGG	7.4 ± 3.4	N.D.
OT3-2	<u>A</u> GTGAGTGAGTGTGT <u>T</u> GTGGGG	24.3 ± 9.2	9.8 ± 4.2
OT3-4	G <u>C</u> TGAGTGAGTGT <u>A</u> TGCGGTG	20.9 ± 11.8	4.2 ± 1.2
OT3-9	GGTGAGTGAGTGC <u>CG</u> TGCG <u>GG</u> GTG	3.2 ± 0.3	N.D.
OT3-17	<u>G</u> TTGAGTGA <u>A</u> TGTGTGCGGTGAGG	2.9 ± 0.2	N.D.
OT3-18	<u>T</u> GT <u>GG</u> GTGAGTGTGTGCGGTGAGG	13.4 ± 4.2	4.9 ± 0.0
OT3-20	<u>A</u> GA <u>A</u> GATGAGTGTGTGC <u>A</u> TGAGG	16.7 ± 3.5	7.9 ± 2.4

Amounts of sgRNA- and Cas9-expressing plasmids transfected into U2OS.EGFP cells for these assays are shown at the top of each column. (Note that data for 250 ng sgRNA/750 ng Cas9 are the same as those presented in **Table 1**.) Mean indel frequencies were determined using the T7EI assay from replicate samples as described in Methods. OT = Off-target sites, numbered as in **Table 1** and

Supplementary Table 2. N.D. = none detected

Supplementary References:

1. Maeder, M.L. et al. Rapid "open-source" engineering of customized zinc-finger nucleases for highly efficient gene modification. *Mol Cell* **31**, 294-301 (2008).
2. Reyon, D. et al. FLASH assembly of TALENs for high-throughput genome editing. *Nat Biotech* **30**, 460-465 (2012).